# **PCT**

123098

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification <sup>6</sup> :			(1	1) International Publication Number:	WO 99/38970	
C12N 15/12, C07K 14/47 1/68, G01N 33/68	, 16/18, C12Q	A1	(4	3) International Publication Date:	5 August 1999 (05.08.99)	
(21) International Application Nun	nber: PCT/IL	99/000	36	(81) Designated States: AL, AM, AT, BY, CA, CH, CN, CU, CZ, DE		
(22) International Filing Date:	21 January 1999 (	21.01.9	9)	GE, GH, GM, HR, HU, ID, II KR, KZ, LC, LK, LR, LS, L1	, IN, IS, JP, KE, KG, KP,	
(30) Priority Data:				MN, MW, MX, NO, NZ, PL, SI SK SI TI TM TR TT I	PT, RO, RU, SD, SE, SG,	

IL

(71) Applicant (for all designated States except US): DIAGNOSTIC TECHNOLOGIES LTD. [IL/IL]; Senate Building, Technion City, 32000 Haifa (IL).

29 January 1998 (29.01.98)

- (72) Inventors; and
  (75) Inventors/Applicants (for US only): ADMON, Arie [IL/IL];
  Deganiot Street 67, 36054 Kiryat Tivon (IL). PALTIELI,
  Yoav [IL/IL]; Einstein Street 75, 34602 Haifa (IL).
  SLOTKY, Ronit [IL/IL]; Galgal Street 8, 34467 Haifa (IL).
- (74) Agent: REINHOLD COHN AND PARTNERS; P.O. Box 4060, 61040 Tel Aviv (IL).

MANDEL, Silvia [IL/IL]; Lea Street 28/1, 34403 Haifa

SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

#### Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

- (54) Title: PLACENTAL PROTEIN 13
- (57) Abstract

The full amino acid and DNA sequences of placental protein 13 (PP13) are disclosed. Also described are various PP13 derived peptide fragments, and a recombinant method for the production of PP13. PP13 may be used in a screening and a diagnostic method for pregnancy-related complications.

## FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of Americ
CA	Canada	IT	lialy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	u	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

15

20

#### PLACENTAL PROTEIN 13

#### FIELD OF THE INVENTION

The present invention relates to a placental protein and its uses.

#### BACKGROUND OF THE INVENTION

References referred to in the text by a number enclosed by parenthesis are listed at the end of the specification.

The goal of pregnancy management is the delivery of a mature, healthy infant, without encountering complications which can adversely affect the well being of both the mother and the newborn. A significant percentage of pregnancies are affected by various disorders. Among these complications are preterm labor and delivery, intrauterine growth retardation and preeclampsia. These conditions negatively impact the outcome of affected pregnancies, at enormous cost both to the patients as well as to the health system.

Placental Protein 13 (PP13) is a protein which was previously isolated from human placental tissue (U.S. 4,500,451 to Bohn, et al., the contents of which are incorporated herein by reference). The protein was characterized by the following parameters: electrophoretic mobility, isoelectric point, sedimentation coefficient, molecular weight determined by ultracentrifugation, molecular weight determined by SDS-PAGE, extinction coefficient and carbohydrate content. The amino acid composition (residues per 100 residues) was determined but not the amino acid sequence.

PP13 was used to develop an assay for the early stage detection of three specific pregnancy-related disorders: intrauterine growth retardation, preeclampsia

- 2 -

and preterm delivery (U.S. 5,198,366 to Silberman). Both a radioimmunoassay (RIA) and an enzyme-linked immunoassay (ELISA) are disclosed using labeled PP13 and anti PP13 antiserum, respectively. No further properties of PP13 are disclosed in the Silberman patent.

5

15

20

**30** 

#### **BRIEF SUMMARY OF THE INVENTION**

It is an object of the present invention to provide a pure PP13 protein.

It is a further object of the present invention to provide a DNA molecule encoding PP13.

10 It is a still further object of the invention to provide a recombinant method for producing PP13.

Additionally, it is an object of the present invention to provide a diagnostic assay based on PP13 for the early detection of pregnancy complications.

It is another object of the invention to provide immunogenic peptides derived from PP13 which can be used in such a diagnostic assay.

According to one aspect of the present invention, there is provided a protein or polypeptide selected from the group consisting of: (a) Placental Protein 13 (PP13) having the amino acid sequence shown in Fig. 2 (SEQ.ID.NO: 9); (b) a polypeptide having a sequence of amino acids included in PP13 and which binds to antibodies which specifically bind to PP13; (c) a protein or polypeptide of (a) or (b) in which one or more amino acids have been added, deleted or replaced without reducing the ability of the protein or polypeptide to bind antibodies which specifically bind to PP13; and (d) a protein or polypeptide having an amino acid sequence including the amino acid sequence of (a) or (b) or (c).

By another aspect of the present invention, there is provided a DNA molecule encoding the above protein or polypeptide.

According to another aspect of the present invention, there is provided a method of screening for pregnancy-related complications comprising the steps of:
(a) providing a serum sample of a pregnant woman; (b) determining the level of PP13 or a peptide derived therefrom in the serum sample, and (c) comparing the determined level with pre-determined normal levels for women at the same

10

15

20

25

30

gestational age, a deviation between the levels being indicative of a pregnancy-related complication.

By one embodiment of the invention, the determination in step (b) is by means of antibodies, preferably monoclonal antibodies, directed against said proteins or polypeptides.

According to yet another aspect of the present invention, there is provided a recombinant method for the production of PP13 comprising inserting said DNA molecule into an expression vector, inserting the expression vector into a host cell, and incubating the host cell under conditions which permit expression of the inserted vector.

The present invention provides for the first time the full amino acid sequence of PP13, as well as its full cDNA sequence. This information can be utilized in a number of applications. For example, modified PP13 protein homologues and analogues can be produced in which one or more amino acids have been added, deleted or replaced, the modified protein typically retaining 75% homology with PP13. Methods for modifying the amino acid sequence of a protein whose full sequence is known are well known in the art, and include e.g. chemical synthesis, controlled mutagenesis and recombinant methods. Such modified proteins may have superior properties over the natural PP13 in various applications, such as superior immunogenicity or immuno-specificity (e.g. the modified protein may be devoid of immune epitopes common with other proteins) for use in an immunoassay for the early detection of pregnancy-related disorders as described in Silberman.

Furthermore, peptide fragments may be prepared from PP13 and such peptides may be modified as described above with respect to the full protein. These peptides may also be used in various applications. For example, it is well known that immunogenic proteins have specific amino acid sequences or epitopes which induce the immune system to mount an immune response to the protein. The above peptides may be tested for the presence of an epitope of PP13 so as to identify the epitope(s). A peptide containing an epitope may then be used in an immunoassay for pregnancy disorders. A number of PP13-derived peptides are disclosed below.

WO 99/38970

The pure PP13 protein or a derived peptide may be used to prepare antibodies to PP13. Either polyclonal or monoclonal antibodies may be produced by

- 4 -

PCT/IL99/00036

standard methods well known to the skilled artisan.

5

10

15

20

25

**30** 

Both the antibodies as well as the proteins and peptides may be used to prepare diagnostic or screening assays for the detection of pregnancy-related complications such as intrauterine growth retardation, preterm delivery and preeclampsia. Examples of such assays are detailed in Silberman, the contents of which are incorporated herein by reference, and include radioimmunoassays (RIA) and enzyme-linked immunoassays (ELISA). In general, such an assay will include the steps of obtaining a serum sample of a pregnant woman, determining the level of PP13 or of a derived peptide in the serum sample by the immunoassay, and comparing the determined level with pre-determined normal levels for women at the same gestational age. A statistically significant deviation between the levels will be indicative of a pregnancy-related complication.

As mentioned above, the full cDNA of PP13 is disclosed here for the first time. Since the full amino acid sequence of PP13 is also disclosed, various DNA molecules encoding PP13 may be prepared due to the degeneracy of the genetic code. In addition, DNA molecules capable of hybridizing to these DNA molecules under stringent conditions may also be prepared. The DNA molecules may be used in a recombinant method for the production of PP13. Such methods are well known in the art and usually involve inserting the DNA molecule into an expression vector such as a plasmid, phage or viral DNA. The expression vector is then inserted into a compatible host cell such as bacterial cells, or eukaryotic cells such as yeast, plant, mammalian or insect cells. The host cell is incubated under conditions which induce expression of the inserted vector, thereby producing PP13.

For example, the DNA encoding PP13 can be inserted into an expression vector under the control of an inducible promotor such as the LacZ promoter, T7 or T4 polymerase promoter, heatshock promoters, etc. One example of an expression vector is the pQE expression vector (QIAGEN). The pQE vector provides high level expression of proteins containing a 6\*His affinity tag in E. coli. The pQE contains a regulatable promoter consisting of the E. coli phage T5

promoter and two *lac* operator sequences. The vector is then inserted into a competent M15 [PREP4] *E. coli* strain (Villarejo and Zabin, 1974). The M15 host cell contains multiple copies of the plasmid pREPA which carries the *lacI* gene encoding the lac repressor. The host cell is incubated with IPTG which rapidly induces expression of the inserted vector, thereby producing PP13. Many other systems may also be used for PP13 expression, as is well known to the skilled artisan.

A kit for diagnosing pregnancy-related complications may be produced based on the present invention. Such a kit, for example, may comprise the following components: (1) antibodies capable of specifically binding PP-13; (2) labeled PP-13, for example by a radioactive, fluorescent or enzyme marker; (3) PP-13 standard solutions at known concentrations; and (4) means for detecting the signal produced in the assay. Such means could be, for example, antiserum raised against the PP-13-binding antibodies.

15

30

10

#### DETAILED DESCRIPTION OF THE DRAWINGS

The present invention will be better understood from the following detailed description of preferred embodiments, taken in conjunction with the following drawings in which:

Fig. 1 shows a partial nucleotide and deduced amino acid sequence of a cDNA from the Expressed Sequence Tag (EST) database (accession R24614). Regions that are similar to the sequenced peptides are underlined. PP13 derived peptide #3 (Fig 1) was found to share partial identity with this cDNA (red underlined letters), and peptides #4, #5 and #6 are 100% identical to the EST database sequence. The nucleotide sequence of the 390-bp cDNA is shown with a translation of the open reading frame (118 amino acids). A Kozak-like translation initiation sequence containing a presumptive start codon (ATG) at nucleotide 33 is labeled with an asterisk. Nucleotide numbers are shown on the left.

Fig. 2 shows the complete nucleotide and deduced amino acid sequence of the PP13 cDNA clone as obtained from RACE analysis. The nucleotide sequence of the 611-bp cDNA is shown with translation of the open reading frame (139 amino

acids). Regions that are identical to the digested peptide are numbered and underlined. A Kozak-like initiation of translation sequence containing a presumptive start codon (ATG) at nucleotide 41 is signed with asterisk. Nucleotide numbers are shown on the left.

Fig. 3 shows the alignment of amino acid sequence of PP13 and eosinophil lysophospholipase (SEQ. ID. NO: 11). Identical amino acids of PP13 protein and eosinophil lysophospholipase (EPL) are designated by bold. There is about 54% identity between the two proteins.

# 10 DETAILED DESCRIPTION OF A PREFERRED EMBODIMENT MATERIALS AND METHODS

#### Materials

15

Modified trypsin and LysC (sequencing grade) were from Promega. Trifluoroacetic Acid (TFA) and hydrogenated Triton X100 (RTX) were from Sigma. Ammonium Carbonate (AC) was from Riedel-de Haen. Acetonitrile (ACN) was from BioLab. 5' and 3' RACE Systems were from Gibco BRL. pUC57 cloning vector (T-Cloning Kit) was from MBI Fermentas.

#### Sequencing the PP13 protein

The PP13 protein was immuno-affinity purified using rabbit polyclonal 20 antibodies raised against placental proteins and affinity purified on the PP-13 protein. In order to further purify the PP-13 protein and to digest it with proteolytic enzymes, we used the method of Rosenfeld et al. (1992) as follows. The PP-13 protein was separated from other contaminating proteins by resolving it on SDS-PAGE in a mini gel format (10x10cm) followed by fixing the gel and staining 25 the gel with Coomassie brilliant blue. The gel was destained in 40% ethanol + 10% acetic acid. The stained gel band containing the PP-13 protein was cut out with a clean razor blade and washed with 50% acetonitrile (ACN) + 200mM Ammonium Carbonate (AC) in water. This treatment was performed in order to remove as much as possible of the SDS, Coomassie brilliant blue and acetic acid. The washed gel 30 piece was air dried for 30 minutes and rehydrated by adding to it 50-100 µl of 200 mM AC + 1% RTX buffer containing 0.5 µgr modified trypsin or 0.5 µgr of LysC.

10

15

20

25

30

After incubation with gentle shaking at 37°C for 12 hours the proteolytic peptides released from the PP-13 protein were eluted from the gel piece by shaking it twice in  $100 \,\mu l$  of 0.1% TFA + 60% ACN at room temperature for 60 min. The solution was separated from the gel piece by centrifugation and dried down in a Speed-Vac to remove excess ACN. The proteolytic peptides were resolved by Reverse-phase HPLC on a Vydac 1x150 mm, C18, 300 .column with a linear gradient from 4% ACN + 0.1% TFA to 60% ACN + 0.085% TFA at room temperature with a flow rate of 40  $\mu l$ /min. The elution pattern of the peptides was determined by UV absorbance at 214 nm and fractions containing peptides were collected by hand into microfuge tube and stored at -80°C. Some of the fractions containing peptides were sequenced on a Protein-Peptide Sequencer (models 476A and 494A, Perkin Elmer) using the manufacturer's standard Edman chemistry and cycles.

cDNA 3' and 5' ends analysis

In order to isolate the full cDNA sequence of the PP-13 gene, we used a standard method called Rapid Amplification of cDNA Ends (RACE) (2) to extend both the 5' and 3' ends of the known parts of the cDNA to its ends. Generally, the RACE method generates cDNA by using a Polymerase Chain Reaction (PCR) to amplify copies of the region between known segments of the cDNA at specific points in the transcript and its 3' or 5' ends. This was accomplished by making copies of the cDNA between synthetic DNA primers complementary to known segments of the message to primers that anneal to the ends of the cDNA.

For the 3' prime end determination, reverse transcriptase (RT) reaction was carried out using 4 µgr of total placental RNA (prepared by TRI reagent from Molecular Research Center, Inc.) and the 3' end primer: (106ras) 5'- ggc cac gcg tcg act agt act ttt ttt ttt ttt ttt ttt - 3'. This was followed by a PCR reaction between the primers: (107ras for the forward reaction) 5'- ggc cac gcg tcg act agt ac - 3' and the reverse primer (100rs, homologous to peptide # 4) was 5'- ggg ata tgg atg ttg gag gag ac - 3'. The PCR reaction included 2.5 mM MgCl<sub>2</sub>, denaturation at 94°C for 45", primer annealing at 60°C for 45" and primer extension at 72°C for 2 min. for 35 cycles.

For the 5' end determination the RT reaction was carried out with 4 ugr

10

of total placental RNA and a specific 3' primer (101ras): 5'- gtc tcc tcc aac atc cat atc - 3'. The 5' end of the cDNA was extended by adding to it poly-dC using the RACE protocol and reagents (Gibco BRL). This was followed by a PCR reaction using conditions as above and the following primers: a backward primer with the abridged anchor primer (AAP) supplied by Gibco BRL and the forward reaction primer 101rs described above.

The resulting PCR fragments were inserted into the pUC57-T cloning vector (T-Cloning Kit #K1212 MBI Fermentas) and clones containing the insert were selected and sequenced by automated DNA at the Biological Services at the Weizmann Institute, Rehovot, Israel.

#### **RESULTS**

#### Identification of peptides from PP13 Protein

In order to either clone the gene encoding the PP-13 protein or to identify its gene in one of the data banks, it was necessary to obtain the primary amino acid sequence of the PP-13 protein. Since the PP13 protein was blocked at its amino terminus, internal amino acid sequences were obtained after proteolytically digesting the protein into peptide fragments. These peptides were separated and purified by chromatography using reverse-phase HPLC, and some of the resolved peptides were sequenced. The amino acid sequences of the peptides that were successfully sequenced are listed in Table 1.

**Table 1.** Amino acid sequences of PP13 derived peptide fragments obtained after trypsin and LysC digestion as described above.

Peptide number	Amino acid sequence
1. (SEQ.ID.NO: 1)	LPVSLSVG
2. (SEQ.ID.NO: 2)	VIIK
3. (SEQ.ID.NO: 3)	GTPIHSFINDPQLQVDF
4. (SEQ.ID.NO: 4)	EFGIWMLEETTDYVPFE
5. (SEQ.ID.NO: 5)	QFELCIY
6. (SEQ.ID.NO: 6)	VHYNEY
7. (SEQ.ID.NO: 7)	GFVHR

15

20

# Comparing peptides sequence to Data-Banks

DNA and protein data banks available through the Internet were searched for homology to the obtained PP-13 peptides sequences. A cDNA sequence (SEQ.ID.NO: 8) encoding four of the peptides fragments (Fig 2) was identified (EST accession R24614). The fact that homology to more than one peptide sequence was present in the identified cDNA indicates that this cDNA is likely a product of the gene encoding the protein which is the major constituent of the PP13 preparation.

The sequence was found in an EST data bank created by the University 10 of Washington and searched through the National Center for Biotechnology Information (NCBI) using the BLAST search program. The R24614 cDNA contains a Kozak-like translation initiation sequence and a 358 base-pair open reading frame (ORF) encoding a 118 amino acid polypeptide. The calculated molecular weight of the polypeptide encoded by the R24614 open reading frame is 13.9 Kda. Four of the sequenced peptides have homology to parts of the deduced sequence of the large open reading frame of the R24614 cDNA (Fig 1). The obtained amino acid sequence of peptide #3 was found to share partial identity with the EST cDNA and peptides number 4, 5 and 6 were identical to different segments of the ORF in the R24614 sequence.

Since the open reading frame sequence of R24614 obtained from the data bank did not contain the entire coding region of the PP13 protein, it was necessary to obtain the full cDNA sequence.

#### Identification of PP13 complete cDNA sequence

25 In order to obtain full cDNA sequence we used Rapid Amplification of cDNA ends (RACE). Using the RACE method with an internal specific primers homologous to the sequence from the region of peptide 4 previously found (Fig 1). we discovered the 3' and 5' end of PP13 message. The full PP13 amino acid sequence (SEQ.ID.NO: 9) and cDNA (SEQ.ID.NO: 10) are shown in Figure 2.

The full cDNA contains a Kozak-like translation initiation sequence and a 417-bp open reading frame encoding a 139 amino acid polypeptide, with a predicted mass of 15.1 KDa which is about the same size of the molecular weight of the PP13 protein as calculated from its migration in SDS-PAGE. The major open reading frame of the full cDNA sequence contains all of the peptides sequence previously found by Edman sequencing of reverse-phase purified proteolytic peptides (Fig 1).

# Resemblance to other proteins

10 It turned out that the novel gene contains sequence similarity to eosinophil lysophospholipase (3), a protein of known significance in immunity and pregnancy disorders (Fig 3). PP13 and eosinophil lysophospholipase have about 54% amino acid identity and 56% nucleic acid identity. The identity of the two proteins in the regions of the peptides, especially peptides number 4 and 6 is low, so it is clear that these proteins are different, but the homology and identity might suggests they belong to the same protein family.

#### References:

- Rosenfeld et al. (1992) In-Gel digestion of protein for internal sequence
   analysis after one or two dimensional Gel Electrophoresis. Analytical Biochemistry,
   203, 173-175.
  - 2. Frohman, M.A., (1990) PCR Protocols: A Guide to Methods and Applications (Innis, M.A., Gelfand, D.H., Sninsky, J.J., and White, T.J, eds.) p. 28, Academic Press, San Diego.
- Ackerman, S.J., Corrette, S.E., Rosenberg, H.F., Bennett, J.C., Mastrianni, D.M., Nicholson-Weller, A., Weller, P.F., Chin, D.T., and Tener, D.G. (1993) The J. of Immunology, 150, No. 2, pp 456-468.
  - 4. Villarego, M.R. and Zabin, I. (1974) J. Bacteriol., 120, 466-474.

#### CLAIMS:

- 1. A protein or polypeptide selected from the group consisting of:
- (a) Placental Protein 13 (PP13) having the amino acid sequence shown in Fig. 2 (SEQ.ID.NO: 9);
- 5 (b) a polypeptide having a sequence of amino acids included in PP13 and which binds to antibodies which specifically bind to PP13;
  - (c) a protein or polypeptide of (a) or (b) in which one or more amino acids have been added, deleted or replaced without reducing the ability of the protein or polypeptide to bind antibodies which specifically bind to PP13; and
- 10 (d) a protein or polypeptide having an amino acid sequence including the amino acid sequence of (a) or (b) or (c).
  - 2. A peptide corresponding to a sub-sequence of PP13 selected from the following group of peptides:

LPVSLSVG (SEQ.ID.NO: 1)

15 VIIK (SEQ.ID.NO: 2)

GTPIHSFINDPQLQVDF (SEQ.ID.NO: 3)

EFGIWMLEETTDYVPFE (SEQ.ID.NO: 4)

QFELCIY (SEQ.ID.NO: 5)

VHYNEY (SEQ.ID.NO: 6)

**20** G F V H R (SEQ.ID.NO: 7).

25

- 3. A DNA molecule encoding the protein or polypeptide of Claim 1.
- 4. A DNA molecule according to Claim 3 having the nucleotide sequence appearing in Fig. 2 (SEQ.ID.NO: 10).
- 5. A DNA molecule capable of hybridizing to the DNA molecule of either of Claims 3 or 4 under stringent conditions.
- 6. Antibodies capable of specifically binding the protein or polypeptide of Claim 1.
- 7. A method of screening for pregnancy-related complications comprising the steps of:
- 30 (a) providing a serum sample of a pregnant woman;

20

- (b) determining the level of the protein of Claim 1(a) or the polypeptide of Claim 1(b) in the serum sample; and
- (c) comparing said determined level with pre-determined normal levels for women at the same gestational age, a deviation between the levels being indicative of a pregnancy-related complication.
- 8. A method according to Claim 7 wherein the determination of step (b) is by means of antibodies capable of specifically binding the protein or polypeptide of Claim 1.
- 9. A method according to Claim 7 wherein said pregnancy-related complication is selected from the group consisting of intrauterine growth retardation, preterm delivery and preclampsia.
  - 10. A recombinant method for the production of PP13 comprising:
    - (a) inserting a DNA molecule according to Claim 3 into an expression vector;
- 15 (b) inserting said expression vector into a host cell; and
  - (c) incubating said host cell under conditions which permit expression of the inserted vector, thereby producing PP13.
  - 11. A kit for diagnosing pregnancy-related complications comprising
    - (a) antibodies capable of specifically binding the protein or polypeptide of Claim 1;
  - (b) labeled PP-13; and
    - (c) PP-13 standard solutions.

#3

M S S L P L Q V D

caattetgaaggtegecaagaaggagagaacaATGTCTTCTTTACCCCTGCAGGTGGAT

E Y T D M D E D S D I A F R F R V H F G 60 TTCTACACTGACATGGATCAGGATTCAGATATTGCCTTCCGTTTCCGAGTGCACTTTGGC

H H V V M N R R E F G I W M L E E T T D

120 AATCATGTGGTCATGAACAGGCGTGAGTTTGGGATGTTGGAGGAGACAACAGAC

Y V P F E D G K Q F E L C I Y V H Y N E

180 TACGTGCCCTTTGAGGATGGCAAACAATTTGAGCTGTGCATCTACGTACATTACAATGAG

#5

Y E I K V N G H T H L R L C P I E S R H
240 TATGAGATAAAGGTCAATGGGCATACGCATTTACGGCTTTGTCCCATCGAATCCCGNCAT

H L L K M G A S V R G D I F P G P S V \_C 300 CATTTETTGAAGATGGGTGCAAGTGTCCGAGGAGATATCTTCCCTGGACCNTCAGTGTGT

V L Q F ? G E M I H 360 GTCTTGCAATTTNAGGGGGAGATGATCCACA

FIG. 1

5' M S S L P V P 1 actggactca attctgaagg tcgccaagaa agaaaaaaca ATGTCTTCTT TACCCGTGCC #2 #3 PVSLSVGSCV IIK GT'PI 61 ATACAAACTG CCTGTGTCTT TGTCTGTTGG TTCCTGCGTG ATAATCAAAG GGACACCAAT HSFIND PQLQ VDF Y TD M D E D 121 CCACTCTTTT ATCAATGACC CACAGCTGCA GGTGGATTTC TACACTGACA TGGATGAGGA SDI'AFR FRVH FGN HVV MNRR 181 TTCAGATATT GCCTTCCGTT TCCGAGTGCA CTTTGGCAAT CATGTGGTCA TGAACAGGCG EFGIWM LEETTDY VPF EDGE 241 TGAGTTTGGG ATATGGATGT TGGAGGAGAC AACAGACTAC GTGCCCTTTG AGGATGGCAA #6 Q F E L C I Y V H Y N E Y E I K V N G I 301 ACAATTTGAG CTGTGCATCT ACGTACATTA CAATGAGTAT GAGATAAAGG TCAATGGCAT RIY GFV HRIP PSF V K M V Q V S 361 ACGCATTTAC GGCTTTGTCC ATCGAATCCC GCCATCATTT GTGAAGATGG TGCAAGTGTC RDISLTSVCVCH 421 GAGAGATATC TCCCTGACCT CAGTGTGTGT CTGCAATtga gggagatgat cacactcctc 481 attgttgagg aaatccctct ttctacctga ccatgggatt cccagaacct gctaacagaa 541 taatecetge teacatttte cectacaett tgteattaaa acageaegaa aacteaaaaa 601 aaaaaaaaaa

FIG. 2

PP13	Ţ	W22T5A51VT5A2T2AG2CATIVG15IH2LIND5ÖFÖADLJ.DWDED
EPL	1	MSLLPVPYTEAASLSTGSTVTIKGRPLVCFLNEPYLQVDFHTEMKEE
PP13	48	SDIAFRFRVHFGNHVVMNRREFGIWMLEETTDYVPFEDGKQFELCIY
		111 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
EPT.	48	SDIVFHFQVCFGRRVVMNSREYGAWKQQVESKNMPFQDGQEFELSIS
	40	ODIA LITTO A CLOSSOL ALMONE LOSMANDO A ESCAPITA L'ODOCE LE L'OLO
:_		
PP13	95	VHYNEYEIKVNGIRIYGFVHRIPPSFVKMVQVSRDISLTSVCVCN
		1 111 1 111 1 111 11 11 11 1
EPL	95	VLPDKYQVMVNGQSSYTFDHRIKPEAVKMVOVWRDISLTKFNVSYT,KR

FIG. 3

#### SEQUENCE LISTING

```
<110> DIAGNOSTIC TECHNOLOGIES LTD.
<120> Placental protein 13
<130> Diagnostic Technologies Ltd.
<140>
<141>
<150> 123098
<151> 1998-01-29
<160> 11
<170> PatentIn Ver. 2.0
<210> 1
<211> 8
<212> PRT
<213> Human placental tissue
<220>
<221> PEPTIDE
<222> (1)..(8)
<223> PP-13 derived (Page 8)
<400> 1
Leu Pro Val Ser Leu Ser Val Gly
  1
<210> 2
<211> 4
<212> PRT
<213> Human placental tissue
<220>
<221> PEPTIDE
<222> (1)..(4)
<223> PP-13 derived (Page 8)
<400> 2
Val Ile Ile Lys
  1
```

```
<210> 3
<211> 17
<212> PRT
<213> Human placental tissue
<220>
<221> PEPTIDE
<222> (1)..(17)
<223> PP-13 derived (Page 8)
<400> 3
Gly Thr Pro Ile His Ser Phe Ile Asn Asp Pro Gln Leu Gln Val Asp
                                     10
                                                          15
Phe
<210> 4
<211> 17
<212> PRT
<213> Human placental tissue
<220>
<221> PEPTIDE
<222> (1)..(17)
<223> PP-13 derived (Page 8)
<400> 4
Glu Phe Gly Ile Trp Met Leu Glu Glu Thr Thr Asp Tyr Val Pro Phe
                                     10
                                                         15
Glu
<210> 5
<211> 7
<212> PRT
<213> Human placental tissue
<220>
<221> PEPTIDE
<222> (1)..(7)
<223> PP-13 derived (Page 8)
<400> 5
Gln Phe Glu Leu Cys Ile Tyr
```

```
1
 <210> 6
 <211> 6
 <212> PRT
 <213> Human placental tissue
 <220>
 <221> PEPTIDE
<222> (1)..(6)
<223> PP-13 derived (Page 8)
<400> 6
Val His Tyr Asn Glu Tyr
  1
<210> 7
<211> 5
<212> PRT
<213> Human placental tissue
<220>
<221> PEPTIDE
<222> (1)..(5)
<223> PP-13 derived (Page 8)
<400> 7
Gly Phe Val His Arg
 1
<210> 8
<211> 611
<212> DNA
<213> Human placental tissue
<220>
<221> gene
<222> (1)..(611)
<223> PP-13 clone R24614 (Fig. 2)
<400> 8
actggactca attctgaagg tcgccaagaa agaaaaaaca atgtcttctt tacccgtgcc 60
atacaaactg cctgtgtctt tgtctgttgg ttcctgcgtg ataatcaaag ggacaccaat 120
ccactctttt atcaatgacc cacagctgca ggtggatttc tacactgaca tggatgagga 180
ttcagatatt gccttccgtt tccgagtgca ctttggcaat catgtggtca tgaacaggcg 240
```

tgagtttggg atatggatgt tggaggagac aacagactac gtgccctttg aggatggcaa 300 acaatttgag ctgtgcatct acgtacatta caatgagtat gagataaagg tcaatggcat 360 acgcatttac ggctttgtcc atcgaatccc gccatcattt gtgaagatgg tgcaagtgtc 420 gagagatatc tecetgaeet cagtgtgtgt etgeaattga gggagatgat cacacteete 480 attgttgagg aaatccctct ttctacctga ccatgggatt cccagaacct gctaacagaa 540 taatccctgc tcacattttc ccctacactt tgtcattaaa acagcacgaa aactcaaaaa 600 aaaaaaaaa a <210> 9 <211> 139 <212> PRT <213> Human placental tissue <220> <221> PEPTIDE <222> (1)..(139) <223> PP-13 (Fig.2) <400> 9 Met Ser Ser Leu Pro Val Pro Tyr Lys Leu Pro Val Ser Leu Ser Val 10 Gly Ser Cys Val Ile Ile Lys Gly Thr Pro Ile His Ser Phe Ile Asn 20 25 Asp Pro Gln Leu Gln Val Asp Phe Tyr Thr Asp Met Asp Glu Asp Ser 35 40 Asp Ile Ala Phe Arg Phe Arg Val His Phe Gly Asn His Val Val Met 50 55 60 Asn Arg Arg Glu Phe Gly Ile Trp Met Leu Glu Glu Thr Thr Asp Tyr 65 75 Val Pro Phe Glu Asp Gly Lys Gln Phe Glu Leu Cys Ile Tyr Val His 90 Tyr Asn Glu Tyr Glu Ile Lys Val Asn Gly Ile Arg Ile Tyr Gly Phe 100 105 Val His Arg Ile Pro Pro Ser Phe Val Lys Met Val Gln Val Ser Arg 115 120 125 Asp Ile Ser Leu Thr Ser Val Cys Val Cys Asn

<210> 10

130

135

```
<211> 417
 <212> DNA
 <213> Human placental tissue
 <220>
 <221> gene
 <222> (1)..(417)
 <223> PP-13 (Fig. 2)
 <400> 10
atgtcttctt tacccgtgcc atacaaactg cctgtgtctt tgtctgttgg ttcctgcgtg 60
ataatcaaag ggacaccaat ccactctttt atcaatgacc cacagctgca ggtggatttc 120
tacactgaca tggatgagga ttcagatatt gccttccgtt tccgagtgca ctttggcaat 180
catgtggtca tgaacaggcg tgagtttggg atatggatgt tggaggagac aacagactac 240
gtgccctttg aggatggcaa acaatttgag ctgtgcatct acgtacatta caatgagtat 300
gagataaagg tcaatggcat acgcatttac ggctttgtcc atcgaatccc gccatcattt 360
gtgaagatgg tgcaagtgtc gagagatatc tccctgacct cagtgtgtgt ctgcaat
<210> 11
<211> 142
<212> PRT
<213> Human white blood cells
<220>
<221> PEPTIDE
<222> (1)..(142)
<223> Eosinophil Lysophospholipase (Fig. 3)
<400> 11
Met Ser Leu Leu Pro Val Pro Tyr Thr Glu Ala Ala Ser Leu Ser Thr
                                      10
Gly Ser Thr Val Thr Ile Lys Gly Arg Pro Leu Val Cys Phe Leu Asn
             20
                                 25
Glu Pro Tyr Leu Gln Val Asp Phe His Thr Glu Met Lys Glu Glu Ser
         35
                             40
Asp Ile Val Phe His Phe Gln Val Cys Phe Gly Arg Arg Val Val Met
     50
Asn Ser Arg Glu Tyr Gly Ala Trp Lys Gln Gln Val Glu Ser Lys Asn
 65
                     70
                                          75
Met Pro Phe Gln Asp Gly Gln Glu Phe Glu Leu Ser Ile Ser Val Leu
                 85
                                     90
Pro Asp Lys Tyr Gln Val Met Val Asn Gly Gln Ser Ser Tyr Thr Phe
```

100 105 110

Asp His Arg Ile Lys Pro Glu Ala Val Lys Met Val Gln Val Trp Arg

Asp Ile Ser Leu Thr Lys Phe Asn Val Ser Tyr Leu Lys Arg 130 135 140

# INTERNATIONAL SEARCH REPORT

Inter onal Application No

			PCI/IL 9	9/00036
A. CLASS IPC 6	IFICATION OF SUBJECT MATTER C12N15/12 C07K14/47 C07K16/	/18 C12Q1/6	8 G01	N33/68
According t	to International Patent Classification (IPC) or to both national classif	ication and IPC		
	SEARCHED			
IPC 6	ocumentation searched (classification system followed by classifica CO7K C12N C12Q			
	tion searched other than minimum documentation to the extent that			
Electronic	tata base consulted during the international search (name of data b	ase and, where practical,	search terms use	d)
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of the re	elevant passages		Relevant to claim No.
X	EP 0 101 603 A (BEHRINGWERKE AG) 29 February 1984 cited in the application see the whole document			1-11
Υ	US 5 198 366 A (SILBERMAN MICHAE 30 March 1993 cited in the application see the whole document	L)		1-11
Y	EP 0 283 606 A (TECHNION RES & D FOUNDATION ;SILBERMAN MICHAEL (I 28 September 1988 see the whole document			1-11
		-/		
X Furth	er documents are listed in the continuation of box C.	X Patent family m	embers are listed	in annex.
* Special cat	egories of cited documents:	*T* later degrament public	had alter the inte	and and title and the
conside	nt defining the general state of the art which is not ared to be of particular relevance ocument but published on or after the International	T later document publis or priority date and r cited to understand invention  "X" document of particula	not in conflict with the principle or th i	the application but sory underlying the
"L" documer which is citation	nt which may throw doubts on priority ctaim(s) or s cited to establish the publication date of another or other special reason (as specified)	cannot be considere involve an inventive "Y" document of particula	d novel or cannot step when the do r relevance; the o	t be considered to current is taken alone staimed invention
"O" docume other m "P" documen	nt referring to an oral disclosure, use, exhibition or neans nt published prior to the international filing date but	document is combin	ed with one or mo	ventive step when the ore other such docu- us to a person skilled
tater th	an the priority date claimed ctual completion of the international search	"&" document member of Date of mailing of the		<u> </u>
25	5 May 1999	04/06/19	99	
Name and m	ailing address of the ISA  European Patent Office, P.B. 5818 Patentlaan 2  NL - 2280 HV Rijswijk  Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax (-31-70) 340-3015	Authorized officer		

# INTERNATIONAL SEARCH REPORT

Inte onal Application No PCT/IL 99/00036

	<del></del>	PC1/1L 99/00036
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
(	BOHN H ET AL: "PURIFICATION AND CHARACTERIZATION OF 2 NEW SOLUBLE PLACENTAL TISSUE PROTEINS PP - 13 AND PP-17."  ONCODEV BIOL MED, (1983) 4 (5), 343-350. CODEN: OBIMD4. ISSN: 0167-1618., XP002103554 see the whole document	1-11
	THAN, GABOR ET AL: "Placental protein (PP5, PP10, PP12, PP13, PP17) levels in sera and in amniotic fluid during healthy pregnancy" MAGY. NOORV. LAPJA (1986), 49(1), 11-15 CODEN: MNLAA8;ISSN: 0025-021X,1986, XP002103555 see the whole document	1-11

# INTERNATIONAL SEARCH REPORT

domation on patent family members

Inter anal Application No
PCT/IL 99/00036

cited	atent document d in search report		Publication date		Patent family member(s)	Publication date
EP	0101603	A	29-02-1984	DE	3230996 A -	23-02-1984
				AT.	38521 T	15-11-1988
				AU	555699 B	02-10-1986
				AU	1816783 A	23-02-1984
				CA	1213213 A	28-10-1986
				DE	3378414 A	15-12-1988
				JP	1693459 C	17-09-1992
				JP	3054680 B	20-08-1991
				JP	59059621 A	05-04-1984
				US	4500451 A	19-02-1985
US	5198366	Α	30-03-1993	CA	12 <b>95</b> 939 A	18-02-1992
ЕР	0283606	Α	28-09-1988	AT	78342 T	15-08-1992
				AU	599639 B	26-07-1990
				AU	7059187 A	22-09-1988
				ZA	8702145 A	26-09-1988